Claims

 A method for preparing closed bacterial ghosts, comprising bringing bacterial ghosts into contact with carrier materials under conditions under which closure of the bacterial ghosts takes place,

characterized in that

the fusion is mediated by way of specific interactions between the partners of a bioaffinity binding pair, which partners are anchored on the ghosts and/or the carrier materials.

The method as claimed in claim 1, characterized in that

the partners of the bioaffinity binding pair are selected from the group consisting of biotin or biotin analogues/streptavidin or avidin, hapten/antibodies or antibody fragments, saccharide/lectin and ligand/receptor.

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3. The method as claimed in claim 2, characterized in that the bioaffinity binding pair employed is biotin/streptavidin.

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4. The method as claimed in one of claims 1 to 3, characterized in that at least one partner of the bioaffinity binding pair is immobilized on the membrane of the bacterial ghosts and on the carrier material.

5. The method as claimed in claim 4, characterized in that

a first partner (P1) of the bioaffinity binding pair is immobilized on the membrane of the bacterial ghosts and a second partner (P2) of the bioaffinity binding pair is immobilized on the carrier material and the closure takes place by way of a P1-P2 interaction.

 The method as claimed in claim 4, characterized in that

a first partner (P1) of the bioaffinity binding pair is immobilized on the membrane of the bacterial ghosts and the carrier material and a second partner (P2) of the bioaffinity binding pair is present in free form and the closure takes place by way of a P1-P2-P1 interaction.

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- 7. The method as claimed in one of the preceding claims, characterized in that the ghosts are derived from Gram-negative bacteria.
- 15 8. The method as claimed in one of the preceding claims, characterized in that the ghosts are derived from recombinant bacteria containing heterologous membrane polypeptides.
- 20 9. The method as claimed in one of the preceding claims, characterized in that the carrier material employed is lipid vesicles.
 - 10. The method as claimed in claim 9,

25 characterized in that

the lipid vesicles employed are vesicles from homogenized cells, in particular bacterial cells, liposomes or membrane-enveloped viruses.

- 30 11. The method as claimed in claim 9 or 10, furthermore comprising an at least partial fusion of the membrane of the bacterial ghosts and the membrane of the lipid vesicles.
- 35 12. The method as claimed in one of the preceding claims, further comprising the packing of active compounds into the bacterial ghosts.
 - 13. The method as claimed in claim 12,

characterized in that

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the active compounds are selected from genetic material, cell components, substances, labeling substances, agriculturally active substances, dyes and combinations thereof.

- 14. A closed bacterial ghost which can be obtained by the method as claimed in one of claims 1 to 13, with the closure being mediated by way of specific interactions between partners of a bioaffinity binding pair.
- 15. The closed bacterial ghost as claimed in claim 14, characterized in that
- it comprises a membrane which is at least partially intact.
 - 16. The closed bacterial ghost as claimed in claim 14 or 15,
- 20 characterized in that

it comprises at least one encapsulated active compound.

- 17. The use of closed bacterial ghosts as claimed in one of claims 14 to 16 in medicine.
 - 18. The use of closed bacterial ghosts as claimed in one of claims 14 to 16 in the agricultural sphere.
- 30 19. The use of closed bacterial ghosts as claimed in one of claims 14 to 16 in biotechnology.